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NOVEL CONCENTRATED INVERSE LATEX, PRODUCTION METHOD AND USE THEREOF IN INDUSTRY

The present patent application relates to thickening water-in-oil inverse latexes, to their process of preparation and to their application as thickener and/or emulsifier in industrial products, in care products for the skin and hair or for the manufacture of cosmetic, dermopharmaceutical or pharmaceutical preparations.

Inverse latexes of polymers of partially or completely salified 2-methyl-2-[(1-oxo-2-propenyl)amino]-1-propane-sulfonic acid (also known as 2-acrylamido-2-methyl-propanesulfonic acid, ATBS or AMPS) and their use in cosmetics and/or pharmaceuticals have formed the subject of numerous patent applications. However, the presence of large amounts of water and oil represents a not insignificant disadvantage in terms of volume, of cost and sometimes of increased risk and/or of toxic effects.

Solutions have thus been developed in order to increase the concentration of polymers in the final latexes, for example by subjecting the reaction medium at the end of the polymerization to a vacuum distillation stage in order to remove a more or less large portion of water and oil. However, this distillation is problematic to carry out as it often destabilizes the reverse latex, which destabilization has to be countered by the prior addition of stabilizing agents. European applications EP 0 161 038 and EP 0 126 528 and British patent application GB 1 482 515 disclose such a use of stabilizing polymers.

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However, these comprise alcohols or glycols which produce environmental problems. Furthermore, the reaction medium sometimes sets solid during the

distillation stage without this phenomenon really ever explained but having the having been consequence of the loss of the batch of reverse latex and laborious and expensive cleaning of the reactor. the distillation Finally, even when takes correctly, the reverse latexes obtained often invert difficulty, they have a high viscosity sometimes internally exhibit microgels. These disadvantages thus forbid them from being used in the manufacture of cosmetic formulations and/or textile 10 printing formulations.

For these reasons, the Applicant Company has endeavored to develop concentrated inverse latexes, that is to say comprising at least 50% by weight of polymer and less than 5% by weight of water, which are devoid of such disadvantages.

According to a first aspect, a subject matter of the invention is a composition in the form of an inverse latex comprising:

- a) from 50% by weight to 80% by weight of at least one linear, branched or crosslinked organic polymer (P),
 - b) from 5% by weight to 10% by weight of an emulsifying system (S_1) of water-in-oil (W/O) type,

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- c) from 5% by weight to 45% by weight of at least one oil, and
- d) from 0% to 5% by weight of water.

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The polymer (P) present in the composition which is a subject matter of the invention can be a homopolymer or a polymer formed from several different types of

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monomers. It is mainly a copolymer, a terpolymer or a tetrapolymer.

The composition as defined above comprises either a single polymer (P) or a blend of different polymers (P).

According to a first specific aspect of the present invention, the polymer (P) is

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- <u>either</u> a homopolymer of a monomer chosen <u>either</u> from those having a partially or completely salified strong acid functional group <u>or</u> from those having a partially or completely salified weak acid functional group <u>or</u> from cationic monomers,
- <u>or</u> a copolymer in which each of the monomers is chosen, independently of one another, <u>either</u> from those having a partially or completely salified strong acid functional group <u>or</u> from those having a partially or completely salified weak acid functional group <u>or</u> from neutral monomers <u>or</u> from cationic monomers,

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- <u>or</u> a terpolymer in which each of the monomers is chosen, independently of one another, <u>either</u> from those having a partially or completely salified strong acid functional group <u>or</u> from those having a partially or completely salified weak acid functional group <u>or</u> from neutral monomers <u>or</u> from cationic monomers,
- <u>or</u> a tetrapolymer in which each of the monomers is chosen, independently of one another, <u>either</u> from those having a partially or completely salified strong acid functional group <u>or</u> from those having a partially or completely salified weak acid

functional group <u>or</u> from neutral monomers <u>or</u> from cationic monomers.

In the composition as defined above, the emulsifying system (S_1) of water-in-oil (W/O) type is composed either of a single surfactant or of a mixture surfactants, provided that said mixture has an HLB value which is sufficiently low to bring about waterin-oil emulsions. Examples of emulsifying agent water-in-oil type include sorbitan esters, such 10 sorbitan oleate, such as that sold by Seppic under the name Montane™ 80, sorbitan isostearate, such as that sold by Seppic under the name Montane TM 70, or sorbitan sesquioleate, such as that sold by Seppic under the name MontaneTM 83. Additional emulsifying agents 15 water-in-oil type are some polyethoxylated sorbitan for example pentaethoxylated sorbitan monooleate, such as that sold by Seppic under the name Montanox[™] 81, or pentaethoxylated sorbitan isostearate, such as that sold under the name Montanox TM 71 by 20 Seppic. Further emulsifying agents of water-in-oil type are diethoxylated oleocetyl alcohol, such as that sold Simulsol™ the name OC 72 by tetraethoxylated lauryl acrylate, such as that sold under the name BlemmerTM ALE 200, or polyesters with a 25 molecular weight of between 1000 and 3000 produced from condensation between a poly(isobutenylsuccinic acid) or its anhydride and a polyethylene glycol, such as Hypermer[™] 2296, sold by Unichema, or, finally, block copolymers with a molecular weight of between 2500 and 30 3500, such as Hypermer[™] B246, sold by Unichema, or Simaline[™] IE 200, sold by Seppic.

The term "branched polymer" denotes, for (P), a nonlinear polymer which has pendent chains so as to obtain, when this polymer is dissolved in water, a high state of entanglement resulting in very high viscosities at low gradient.

The term "crosslinked polymer" denotes, for (P), a nonlinear polymer existing in the form of a three-dimensional network which is insoluble in water but which is swellable in water and which thus results in a chemical gel being obtained.

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The composition according to the invention can comprise linear units, crosslinked units and/or branched units.

When the polymer (P) is crosslinked, it is crosslinked 10 more particularly with a diethylene or polyethylene in the molar proportion, expressed with compound respect to the monomers employed, of 0.005% to 1%, preferably of 0.01% to 0.2% and more particularly of 15 0.01% to 0.1%. Preferably, the crosslinking and/or the branching agent is chosen from ethylene glycol dimethacrylate, diethylene glycol diacrylate, sodium diallyloxyacetate, ethylene glycol diacrylate, diallylurea, triallylamine, trimethylolpropane triacrylate or methylenebis(acrylamide). 20

strong acid functional group of the monomers is in particular the sulfonic acid comprising it functional group or the phosphonic acid functional group. Said monomers are, for example, partially or completely salified styrenesulfonic acid preferably, partially or completely salified 2-methyl-2-[(1-oxo-2-propenyl)amino]-1-propanesulfonic (also known as 2-acrylamido-2-methylpropanesulfonic acid).

The weak acid functional group of the monomers comprising it is in particular the partially salified carboxylic acid functional group. Said monomers can, for example, be partially or completely salified acrylic acid, methacrylic acid, itaconic acid, maleic acid or 3-methyl-3-[(1-oxo-2-propenyl)amino]butanoic acid.

monomers are chosen in particular from The neutral methacrylamide, diacetone acrylamide, acrylamide, N-isopropylacrylamide, N-[2dimethylacrylamide, hydroxy-1,1-bis(hydroxymethyl)ethyl]propenamide tris(hydroxymethyl)acrylamidomethane orN-[tris(hydroxymethyl) methyl] acrylamide, also known as 2-hydroxyethyl acrylate, 2,3-dihydroxypropyl acrylate, methacrylate, 2,3-dihydroxypropyl 2-hydroxyethyl with methacrylate, an ethoxylated derivative molecular weight of between 400 and 1000 of each of these esters, or vinylpyrrolidone.

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The cationic monomers are chosen in particular from quaternary ammonium derivatives. Said monomers can, for 15 example, be 2, N, N, N-tetramethyl-2-[(1-oxo-2-2, N, N-trimethyl-2-[(1propenyl) amino] propanammonium, oxo-2-propenyl) amino] propanammonium, N,N,N-trimethyl-2-[(1-oxo-2-propenyl)oxy]ethanammonium, N,N,N-trimethyl-3-[(1-oxo-2-propenyl)oxy]propanammonium, N,N,N-trimethyl-20 2-[(1-oxo-2-propenyl)amino]propanammonium diallylordimethylammonium salts. The term "salt" is understood to mean more particularly the chlorides, the bromides or the iodides of said ammonium salts.

25 For the monomers comprising a strong acid functional group or comprising a weak acid functional group, the term "salified" means the alkali metal salts, such as the sodium or potassium salts, or the salts of nitrogenous bases, such as the ammonium salt or the 30 monoethanolamine (HO-CH₂-CH₂-NH₄⁺) salt.

According to a second specific aspect of the present invention, the polymer (P) is chosen from:

35 - crosslinked copolymers of acrylic acid, partially salified in the sodium salt or ammonium salt form, and of acrylamide; - 7 -

- crosslinked copolymers of 2-methyl-2-[(1-oxo-2-propenyl)amino]-1-propanesulfonic acid, partially salified in the sodium salt form, and of acrylamide;

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- crosslinked copolymers of 2-methyl-2-[(1-oxo-2-propenyl)amino]-1-propanesulfonic acid and of acrylic acid, which are partially salified in the sodium salt form;

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- crosslinked copolymers of 2-methyl-2-[(1-oxo-2-propenyl)amino]-1-propanesulfonic acid, partially salified in the sodium salt form, and of 2-hydroxyethyl acrylate;

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- crosslinked copolymers of acrylamide and of N,N,Ntrimethyl-3-(1-oxo-2-propenyl)propanammonium;
- crosslinked homopolymers of 2-methyl-2-[(1-oxo-2-20 propenyl)amino]-1-propanesulfonic acid, partially salified in the sodium salt form;
- crosslinked homopolymers of acrylic acid,
 partially salified in the ammonium salt or
 monoethanolamine salt form;
 - terpolymers of acrylamide, of N,N,N-trimethyl-3-(1-oxo-2-propenyl)propanammonium and of [tris(hydroxymethyl)aminomethyl]acrylamide;

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- crosslinked terpolymers of acrylamide, of 2-methyl-2-[(1-oxo-2-propenyl)amino]-1-propane-sulfonic acid and of acrylic acid, which are partially salified in the sodium salt form;

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- terpolymers of 2-methyl-2-[(1-oxo-2-propenyl)amino]-1-propanesulfonic acid, partially salified in the sodium salt form, of acrylamide and of vinylpyrrolidone. According to a third specific form of the present invention, the composition as defined above comprises at least 60% by weight and at most 70% by weight of polymer (P).

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According to a fourth specific form of the present invention, the composition as defined above additionally comprises up to 5% of its weight of an emulsifying system (S_2) of oil-in-water (O/W) type.

"emulsifying agent of oil-in-water type" term denotes emulsifying agents having an HLB value which is sufficiently high to provide oil-in-water emulsions, such as ethoxylated sorbitan esters, for example the sorbitan oleate polyethoxylated with 20 mol of ethylene oxide sold by Seppic under the name of Montanox M 80, the sorbitan laurate polyethoxylated with 20 mol ethylene oxide sold by Seppic under the name MontanoxTM 20, the castor oil polyethoxylated with 40 mol of ethylene oxide sold under the name $Simulsol^{TM}$ OL50, the decaethoxylated oleodecyl alcohol sold by Seppic under the name of Simulsol™ OC 710, the heptaethoxylated lauryl alcohol sold under the name of SimulsolTM P7, the decaethoxylated nonylphenol SynperonicTM the name of NP-10 or the polyethoxylated sorbitan hexaoleates sold by Atlas under the names of G-1086 and G-1096.

In the composition which is a subject matter of the 30 present invention, the oil phase is composed either of mineral comprising commercial oil saturated hydrocarbons, such as paraffins, isoparaffins cycloparaffins, and exhibiting, at ambient temperature, a density between 0.7 and 0.9 and a boiling point of 35 greater than approximately 250°C, such as, for example, Marcol[™]52, sold by Exxon Chemical, or of a vegetable such as squalane of vegetable origin, or of a synthetic oil, such as hydrogenated polyisobutene or hydrogenated polydecene, or of a mixture of several of these oils. MarcolTM 52 is a commercial oil corresponding to the definition of liquid paraffins of the French Pharmacopeia. It is a white mineral oil in accordance with the FDA 21 CFR 172.878 and CFR 178.3620 (a) regulations and is included in the United States Pharmacopeia, US XXIII (1995), and in the European Pharmacopeia (1993). The composition according to the invention can also comprise various additives, such as complexing agents, chain transfer agents or chain-limiting agents.

According to another aspect of the present invention, a subject matter of the latter is a process for the preparation of the composition as defined above, characterized in that:

a) an aqueous phase (A) comprising the monomers and the optional hydrophilic additives is emulsified in an organic phase (O) comprising the surfactant system (S_1) , a mixture composed of the oil intended to be present in the final composition and of a volatile oil, and the optional hydrophobic additives,

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b) the polymerization reaction is initiated by introduction of an initiator of free radicals into the emulsion formed in a) and then the reaction is allowed to take place, and

- c) the reaction medium resulting from stage b) is concentrated by distillation until said volatile oil has been completely removed.
- The volatile oils appropriate for the implementation of the process as defined above are, for example, light isoparaffins comprising from 8 to 11 carbon atoms, such as, for example, those sold under the names of IsoparTM G, IsoparTM L, IsoparTM H or IsoparTM J.

According to a preferred implementation of the process as defined above, the polymerization reaction is initiated by an oxidation/reduction couple, such as the cumene hydroperoxide/sodium metabisulfite couple, at a temperature of less than or equal to 10°C, and is then carried out either quasiadiabatically up to a temperature of greater than or equal to 40°C, more particularly of greater than or equal to 50°C, or by controlling the change in the temperature.

When stage c) is complete, one or more emulsifying agents of oil-in-water type are introduced, if desired, at a temperature of less than 50°C.

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Another subject matter of the invention is the use of the composition as defined above in preparing a cosmetic, dermopharmaceutical or pharmaceutical topical composition.

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A topical composition according to the invention, intended to be applied to the skin or mucous membranes of man or animals, can consist of a topical emulsion comprising at least one aqueous phase and at least one oil phase. This topical emulsion can be of the oil-in-water type. More particularly, this topical emulsion can consist of a fluid emulsion, such as a milk or a fluid gel. The oil phase of the topical emulsion can consist of a mixture of one or more oils.

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A topical composition according to the invention may be intended for a cosmetic use or may be used to prepare a medicament intended for the treatment of diseases of the skin and mucous membranes. In the latter case, the topical composition then comprises an active principle which can, for example, consist of an antiinflammatory agent, a muscle relaxant, an antifungal or antibacterial.

When the topical composition is used as cosmetic composition intended to be applied to the skin or mucous membranes, it may or may not comprise an active principle, for example a moisturizing agent, a tanning agent, a sunscreen, an agent for combating wrinkles, an agent with a slimming purpose, an agent for combating free radicals, an antiacne agent or antifungal.

A topical composition according to the invention usually comprises between 0.1% and 10% by weight of the thickening agent defined above. The pH of the topical composition is preferably greater than or equal to 5.

The topical composition can additionally comprise compounds conventionally present in compositions of this type, for example fragrances, preservatives, colorants, emollients or surfactants.

According to yet another aspect, the invention relates to the use of the novel thickening agent in accordance with the invention mentioned above for thickening and emulsifying a topical composition comprising at least one aqueous phase.

- The composition according to the invention is an advantageous substitute for those sold under the names of Sepigel™ 305, Sepigel™ 501, Simulgel™ EG, Simulgel™ NS or Simulgel™ 600 by the Applicant Company as it is also very compatible with the other excipients used in the preparation of formulations such as milks, lotions, creams, salts, baths, balms, shampoos or conditioners. They can also be employed with said Sepigel or Simulgel products.
- It is in particular compatible with the concentrates disclosed and claimed in international publications WO 92/06778, WO 95/04592, WO 95/13863, WO 96/37285, WO 98/22207 or WO 98/47610 or in FR 2734 496 and with the surface-active agents disclosed in WO 93/08204.

It is particularly compatible with Montanov[™] 68, Montanov[™] 82, Montanov[™] 202 or Sepiperl[™] N. It can also be used in emulsions of the type of those disclosed and claimed in EP 0 629 396 and in the aqueous dispersions which are cosmetically or physiologically acceptable with an organopolysiloxane compound chosen, for example, from those disclosed in WO 93/05762 or in WO 93/21316.

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also be used to form cosmetically physiologically acceptable aqueous gels at acidic pH, such as those disclosed in WO 93/07856; it can also be used in combination with nonionic celluloses, example to form styling gels, such as those disclosed in EP 0 684 024, or also in combination with esters of fatty acids and of sugar, to form compositions for the the hair or the skin, treatment of such as disclosed in EP 0 603 019, or also in shampoos disclosed conditioners, such as and claimed in WO 92/21316, or, finally, in combination with anionic homopolymer, such as CarbopolTM, to form hair treatment products, such as those disclosed in DE 195 23596, in combination with other thickening or polymers.

The composition according to the invention is also compatible with the active principles, such as, for example, self-tanning agents, such as dihydroxyacetone (DHA), or antiacne agents; they can thus be introduced into self-tanning compositions, such as those claimed in EP 0 715 845, EP 0 604 249, EP 0 576 188 or in WO 93/07902.

35 It is also compatible with the N-acylated derivatives of amino acids, which allows it to be used in soothing compositions, in particular for sensitive skin, such as those disclosed or claimed in WO 92/21318, WO 94/27561 or in WO 98/09611.

When the composition as defined above is intended for the treatment of the hair, it more particularly comprises an inverse latex of cationic polymer which is a subject matter of the present invention.

When the composition as defined above is intended for the treatment of the skin and/or mucous membranes, it more particularly comprises an inverse latex of anionic polymer which is a subject matter of the present invention.

The inverse latexes which are a subject matter of the present invention can be used as thickeners for textile printing pastes.

The aim of the following examples is to illustrate the present invention.

20 EXAMPLE 1: Inverse latex of the AM/APTAC/THAM terpolymer (molar ratio of monomers: 73/20/7) (Cationic thickener - Composition 1)

- a) The following are successively introduced, with25 stirring, into a first beaker:
 - 388.8 g of a commercial 50% by weight acrylamide (AM) solution
 - 206.5 g of a commercial 75% N,N,N-trimethyl-3-[(1-oxo-2-propenyl)propanammonium chloride (APTAC) solution
 - 46 g of [tris(hydroxymethyl)aminomethyl]acrylamide
 (THAM)
 - 0.56 g of commercial 40% sodium diethylenetriaminepentaacetate solution, and
- 35 deionized water, so as to bring the total weight to 813.8 g
 - The pH is adjusted to 5.

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- b) An organic phase is prepared in a second beaker by mixing:
- 137.5 g of Marcol[™] 52
- 186.3 g of Isopar™ G
- 5 25 g of Montane[™] 70 (sorbitan isostearate)
 - 6.2 g of Hypermer[™] 2296
 - 6.0 g of Simaline™ IE 200
 - 6.2 g of tetraethoxylated lauryl acrylate,
 - 125 g of azobis(isobutyronitrile) (AIBN)

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- c) The two phases are subsequently mixed with stirring and subjected to vigorous mechanical stirring so as to create a fine emulsion. This emulsion is subsequently placed in a reactor and nitrogen is sparged therein in order to remove the dissolved oxygen therefrom.
- d) After cooling to approximately 8°C, the polymerization reaction is initiated using the oxidation/reduction couple: cumene hydroperoxide/sodium metabisulfite.
- e) Once the polymerization reaction is complete, the $Isopar^{TM}$ G and virtually all the water are removed by vacuum distillation.
- f) After introduction of 5% of Montanox[™] 20, a cationic thickening inverse latex is obtained which comprises approximately 63% of polymer. The product obtained is devoid of particles and of microgel. It is not very viscous, it has a high thickening power and it readily inverts. Its water content, measured by Karl-Fischer titrimetry, is 3% by weight.

Viscosity measurements (Brookfield RVT viscometer)

	Spindle (S); Speed of	Viscosity in
	rotation of the	mPa·s
	spindle (SR) (in	
	revolutions per	
	minute)	
Inverse latex	S 4; SR: 20	6000
2% by weight	S 6; SR: 5	139 000
aqueous solution		
2% by weight	S 6; SR: 5	12 900
aqueous solution		
+ 0.1% by weight		
of NaCl		

EXAMPLE 2: Inverse latex of the AM/ATBS copolymer (molar ratio: 70/30) crosslinked with MBA (Anionic thickener - Composition 2)

- a) The following are successively introduced, with stirring, into a first reactor:
- 10 245 kg of a commercial 50% by weight acrylamide (AM) solution
 - 308.1 kg of a commercial solution comprising 55% of the sodium salt of 2-acrylamido-2-methylpropanesulfonic acid (ATBS)
- 15 0.066 kg of methylenebis(acrylamide) (MBA)
 - 0.37 kg of a commercial 40% sodium diethylenetriaminepentaacetate solution
 - the pH is adjusted to 5.0 using powdered 2acrylamido-2-methylpropanesulfonic acid
- 20 deionized water, so as to bring the total weight to 564.3 kg.
 - b) An organic phase is prepared in a second reactor by mixing:
- 25 107.6 kg of polyisobutene
 - 74.5 kg of Isopar™ G
 - 14.1 kg of MontaneTM 70

- 2.5 kg of Hypermer[™] 2296
- 4.1 kg of Simaline™ IE 200
- c) The aqueous phase is then introduced into the organic phase with stirring and then the pre-emulsion thus obtained is subjected to vigorous mechanical stirring using a turbine mixer of Silverson type so as to create a fine emulsion while sparging with nitrogen.
- 10 d) After cooling to approximately 8°C, the polymerization reaction is initiated using the oxidation/reduction couple: ammonium persulfate/sodium metabisulfite.
- 15 e) Once the polymerization reaction is complete, the $Isopar^{TM}$ G and virtually all the water are removed by vacuum distillation.
- f) After introduction of 5% of Montanox[™] 20, an anionic thickening inverse latex is obtained which comprises approximately 63% of polymer. The product obtained is not very viscous, it has a high thickening power and it readily becomes inverted. Its water content, measured by Karl-Fischer titrimetry, is 3% by weight.

Viscosity measurements (Brookfield RVT viscometer)

	Spindle (S); Speed of	Viscosity in
	rotation of the	mPa·s
	spindle (SR) (in	
	revolutions per	
	minute)	
Inverse latex	S 4; SR: 20	4000
2% by weight	S 6; SR: 5	135 000
aqueous solution		
2% by weight	S 6; SR: 5	20 800
aqueous solution		
+ 0.1% by weight		

of NaCl

EXAMPLE 3: Inverse latex of the AM/AA copolymer (molar ratio: 25/75) (Anionic thickener - Composition 3)

- 5 a) The following are successively introduced, with stirring, into a first beaker:
 - 106.5 g of a commercial 50% (by weight) acrylamide (AM) solution
 - 162.0 g of glacial acrylic acid (AA)
- 10 98.1 g of a 29.3% by weight aqueous ammonia solution
 - 277 g of methylenebis(acrylamide) (MBA)
 - 0.45 g of a commercial 40% sodium diethylenetriaminepentaacetate solution
- 15 deionized water, up to 680 g
 - b) An organic phase is prepared in a second beaker by mixing:
 - 121 g of Marcol[™] 52
- 20 99 q of Isopar[™] G
 - 17 g of MontaneTM 70
 - 3 kg of HypermerTM 2296
 - 5 g of Simaline[™] IE 200
 - 0.1 q of AIBN

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- c) The aqueous phase is then introduced into the organic phase with stirring and then the pre-emulsion thus obtained is subjected to vigorous mechanical stirring using a turbine mixer of Silverson type so as to create a fine emulsion while sparging with nitrogen.
- d) After cooling to approximately 8°C, the polymerization reaction is initiated using the oxidation/reduction couple: cumene hydroperoxide/sodium metabisulfite.

- e) Once the polymerization reaction is complete, the $Isopar^{TM}$ G and virtually all the water are removed by vacuum distillation.
- 5 f) After introduction of 5% of Montanox[™] 20, an anionic thickening inverse latex is obtained which comprises approximately 63% of polymer. The product obtained is not very viscous, it has a high thickening power and it readily becomes inverted. Its water content, measured by Karl-Fischer titrimetry, is 2.5% by weight.

Viscosity measurements (Brookfield RVT viscometer)

15 A - The viscosities of an aqueous solution comprising 2% by weight of the concentrated inverse latex obtained and of an aqueous solution comprising 2% by weight of said inverse latex and 0.1% by weight of sodium chloride are measured.

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	Spindle (S); Speed of	Viscosity in
	rotation of the	mPa·s
	spindle (SR) (in	
	revolutions per	
	minute)	
Inverse latex	S 4; SR: 20	400
2% by weight	S 6; SR: 5	150 000
aqueous solution		
2% by weight	S 6; SR: 5	72 800
aqueous solution		
+ 0.1% by weight		
of NaCl		

B - a nonconcentrated inverse latex is prepared by carrying out stages a) to d) of the process described in the present example with the same amounts of products.

On conclusion of stage d), 5% of MontanoxTM 20 is added and an inverse latex (Composition III) is obtained which comprises 28% of polymer.

The viscosity of the following solutions is measured:

Composition T at 2% in water: Solution S₁

2% Composition T + 0.1% by weight of NaCl: S₂

Composition 3 at 1% in water: Solution S₃

Composition 3 at 1% in water + 0.1% by weight of NaCl:

Solution S₄

The following results are obtained:

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	Spindle (S); Speed of	Viscosity in
	rotation of the	mPa·s
	spindle (SR) (in	
	revolutions per	
	minute)	
S ₁ (state of the	S 6; SR: 5	47 800
art)		
S ₃ (invention)	S 6; SR: 5	55 400
S ₂ (state of the	S 3; SR: 5	560
art)		
S ₄ (invention)	S 3; SR: 5	1400

The comparison of the results of the salt-comprising solutions (S_2) and (S_4) reveals that the concentrated inverse latex behaves better towards salts than the inverse latex of the state of the art at an equivalent concentration of polymer.

EXAMPLE 4: Inverse latex of the AM/ATBS/AA terpolymer (molar ratio: 65/30/) (Anionic thickener - Composition 4)

- a) The following are successively introduced, with25 stirring, into a first reactor:
 - 227.5 kg of a commercial 50% (by weight) acrylamide (AM) solution

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- 308.1 kg of a commercial solution comprising 55% of the sodium salt of 2-acrylamido-2-methyl-propanesulfonic acid (ATBS)
- 8.8 kg of acrylic acid (AA)
- 5 0.066 kg of methylenebis(acrylamide) (MBA)
 - 0.37 kg of a commercial 40% sodium diethylenetriaminepentaacetate solution
 - the pH is adjusted to 6.2 using sodium hydroxide
 - deionized water, so as to bring the total weight to 564.3 kg.
 - b) An organic phase is prepared in a second reactor by mixing:
 - 107.6 kg of polyisobutene
- 15 74.5 kg of Isopar™ G
 - 14.1 kg of Montane[™] 70
 - 2.5 kg of HypermerTM 2296
 - 4.1 kg of Simaline™ IE 200
- c) The aqueous phase is then introduced into the organic phase with stirring and then the pre-emulsion thus obtained is subjected to vigorous mechanical stirring using a turbine mixer of Silverson type so as to create a fine emulsion while sparging with nitrogen.

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d) - After cooling to approximately 8°C, the polymerization reaction is initiated using the oxidation/reduction couple: ammonium persulfate/sodium metabisulfite.

- e) Once the polymerization reaction is complete, the $Isopar^{TM}$ G and virtually all the water are removed by vacuum distillation.
- 35 f) After introduction of 5% of Montanox[™] 20, an anionic thickening inverse latex is obtained which comprises approximately 63% of polymer. The product obtained is not very viscous, it has a high thickening power and it readily becomes inverted. Its water

content, measured by Karl-Fischer titrimetry, is 3% by weight.

Viscosity measurements (Brookfield RVT viscometer)

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	Spindle (S); Speed of	Viscosity in
	rotation of the	mPa·s
	spindle (SR) (in	
	revolutions per	
	minute)	
Inverse latex	S 4; SR: 20	4000
2% by weight	S 6; SR: 5	90 000
aqueous solution		
2% by weight	S 6; SR: 5	20 000
aqueous solution		
+ 0.1% by weight		
of NaCl		

EXAMPLE 5: Inverse latex of the AM/ATBS/VP terpolymer (molar ratio: 65/25/10) (Anionic thickener - Composition 5)

- a) The following are successively introduced, with stirring, into a first beaker:
- 245.6 g of a commercial 50% (by weight) acrylamide (AM) solution
- 15 279 g of a commercial solution comprising 55% of the sodium salt of 2-acrylamido-2-methyl-propanesulfonic acid (AMPS)
 - 29.6 g of vinylpyrrolidone (VP)
 - 0.082 g of methylenebis(acrylamide) (MBA)
- 20 0.45 g of a commercial 40% sodium diethylenetriaminepentaacetate solution
 - the pH is adjusted to 5.0 using powdered 2-acrylamido-2-methylpropanesulfonic acid
- deionized water, so as to bring the total weight to 644.7 g.

- b) An organic phase is prepared in a second beaker by mixing:
- 110 g of polyisobutene
- 133 g of Isopar™ G
- 5 $13.5 \text{ g of Montane}^{\text{TM}}$ 70
 - 6.5 q of MontanoxTM 71
 - 3.0 g of Hypermer[™] 2296
 - 5.0 g of Simaline[™] IE 200
- 10 c) The aqueous phase is then introduced into the organic phase with stirring and then the pre-emulsion thus obtained is subjected to vigorous mechanical stirring using a turbine mixer of Silverson type so as to create a fine emulsion while sparging with nitrogen.

15

d) - After cooling to approximately 8°C, the polymerization reaction is initiated using the oxidation/reduction couple: ammonium persulfate/sodium metabisulfite.

- e) Once the polymerization reaction is complete, the $Isopar^{TM}$ G and virtually all the water are removed by vacuum distillation.
- 25 f) After introduction of 5% of Montanox[™] 20, an anionic thickening inverse latex is obtained which comprises approximately 63% of polymer. The product obtained is not very viscous, it has a high thickening power and it readily becomes inverted. Its water
- 30 content, measured by Karl-Fischer titrimetry, is 4% by weight.

Viscosity measurements (Brookfield RVT viscometer)

	Spindle (S); Speed of	Viscosity in
	rotation of the	mPa·s
	spindle (SR) (in	
	revolutions per	
	minute)	
Inverse latex	S 4; SR: 20	4000
2% by weight	S 6; SR: 5	75 000
aqueous solution		
2% by weight	S 6; SR: 5	10 000
aqueous solution		
+ 0.1% by weight		
of NaCl		

5 EXAMPLE 6: Inverse latex of the AM/APTAC copolymer (molar ratio: 85/15) (Cationic thickener - Composition)

- a) The following are successively introduced, with stirring, into a first beaker:
- 10 452.6 g of a commercial 50% (by weight) acrylamide (AM) solution
 - 154.9 g of a commercial 75% N,N,N-trimethyl-3-(1-oxo-2-propenyl)propanammonium chloride (APTAC) solution
- 15 0.029 g of methylenebis(acrylamide)
 - 0.56 g of a commercial 40% sodium diethylenetriaminepentaacetate solution
 - the pH is adjusted to 5.0
- the total amount is adjusted to 814 g by addition of the remaining water.
 - b) An organic phase is prepared in a second beaker by mixing:
 - 137.5 g of Marcol[™] 52
- 25 186.5 g of Isopar[™] G
 - 25.1 g of Montane[™] 70 (sorbitan isostearate)
 - 6.3 g of Hypermer[™] 2296

- 6.3 g of ethoxylated lauryl acrylate comprising $4 \text{ mol (Blemmer}^{TM} \text{ ALE 200)}$
- 0.123 g of azobis(isobutyronitrile)
- 5 c) The aqueous phase is then introduced into the organic phase with stirring and then the pre-emulsion thus obtained is subjected to vigorous mechanical stirring using a turbine mixer of Silverson type so as to create a fine emulsion. The combined mixture is also placed under nitrogen sparging.
 - d) The polymerization is then initiated using the oxidizing system; cumene hydroperoxide and ammonium persulfate, and reducing system; sodium metabisulfite.
- 15 e) Once the polymerization reaction is complete, the $Isopar^{TM}$ G and virtually all the water are removed by vacuum distillation.
- f) 5% of MontanoxTM 20 is added, so as to render the latex self-invertible.

The product obtained is not very viscous, it readily becomes inverted and it has a high thickening power. Its water content, measured by Karl-Fischer titrimetry, is 3% by weight.

Viscosity measurements (Brookfield RVT viscometer)

	Spindle (S); Speed of	Viscosity in
	rotation of the	mPa·s
	spindle (SR) (in	
	revolutions per	
	minute)	
Inverse latex	S 4; SR: 20	1300
2% by weight	S 6; SR: 5	80 600
aqueous solution		
2% by weight	S 6; SR: 5	13 000
aqueous solution		

+ 0.1% by weight	
of NaCl	

Examples of cosmetic formulations

Example 7: Care cream

5	Cyclomethicone:		10%
	Compound of example 2:		0.8%
	Montanov TM 68:		4.5%
	Preservative:		0.65%
	Lysine:		0.025%
10	EDTA (disodium salt):		0.05%
	Xanthan gum:		0.2%
	Glycerol:		3%
	Water:	q.s. for	100%

15 Example 8: Care cream

	Cyclomethicone:		10%
	Compound of example 4:		0.8%
	Montanov TM 68:		4.5%
	Perfluoropolymethylisopropyl	ether:	0.5%
20	Preservative:		0.65%
	Lysine:		0.025%
	EDTA (disodium salt):		0.05%
	Pemulen [™] TR:		0.2%
	Glycerol:		3%
25	Water:	q.s. for	100%

Example 9: Aftershave balm

FORMULA

	A	Composition of example 3:		1.5%
30		Water:	q.s. for	100%
	В	Micropearl™ M 100:		5.0%
		Sepicide [™] CI:		0.50%
		Fragrance:		0.20%
35		95° Ethanol:		10.0%

PROCEDURE

B is added to A.

Example 10: Satin emulsion for the body

5 FORMULA

	1 0101	<u>OLLI</u>		
	A	Simulsol [™] 165:		5.0%
		Lanol™ 1688:		8.50%
		Shea butter:		2%
		Liquid paraffin:		6.5%
10		Lanol TM 14M:		3%
		Lanol™ S:		0.6%
	В	Water:		66.2%
15	С	Micropearl TM M 100:		5%
	D	Compound of example 5:		3%
	E	Sepicide [™] CI:		0.3%
20		Sepicide™ HB:		0.5%
		Monteine™ CA:		1%
		Fragrance:		0.20%
		Vitamin E acetate:		0.20%
		Sodium pyrrolidinonecarboxylat	e:	1%
25			(moisturi:	zing agent)

PROCEDURE

C is added to B, B is emulsified in A at 70° C, D is then added at 60° C and then E is added at 30° C.

30

Example 11: Body milk

FORMULA

A	Simulsol™ 165:	5.0%
	Lanol™ 1688:	12.0%
35	Lanol TM 14 M:	2.0%
	Cetyl alcohol:	0.3%
	Schercemol™ OP:	3%

B Water:

q.s. for 100%

	С	Compound of example 4:	0.35%
5	D	Sepicide [™] CI: Sepicide [™] HB: Fragrance:	0.2% 0.5% 0.20%
10	B is	EDURE semulsified in A at approximately 75°C, approximately 60°C and then D is coximately 30°C.	C is added added at
15	Exam FORM A	The state of the s	5.0% 20.0% 1.0%
20	В	Water: q.s. for	2.50%
	D	Compound of example 2: Sepicide [™] CI: Sepicide [™] HB:	0.20%
25	PROC	'EDURE	
	B i	s introduced into A at approximately d at approximately coximately 45°C.	
30	Exam FORM	ple 13: Non-greasy antisun gel	
2.5	A	Compound of example 5: Water:	3.00% 30%
35	В	Sepicide [™] CI: Sepicide [™] HB:	0.20%

0.10%

Fragrance:

C Colorant: q.s. Water: 30%

D MicropearlTM M 100: 3.00% Water: q.s. for 100%

E Silicone oil: 2.0%
Parsol™ MCX: 5.00%

10 PROCEDURE

B is introduced into A, C is added, then D is added and then E is added.

Example 14: Antisun milk

15 FORMULA

A SepiperlTM N: 3.0% Sesame oil: 5.0% ParsolTM MCX: 5.0% λ -Carrageenan: 0.10%

20

5

B Water: q.s. for 100%

C Compound of example 3: 0.80%

25 D Fragrance: q.s. Preservative: q.s.

PROCEDURE

B is emulsified in A at 75°C, then C is added at approximately 60°C, then D is added at approximately 30°C and the pH is adjusted, if necessary.

Example 15: Massage gel

FORMULA

35 A Compound of example 2: 3.5% Water: 20.0%

B Colorant: 2 drops/100 g Water: q.s.

	C	Alcohol:	10%
		Menthol:	0.10%
5	D	Silicone oil:	5.0%
		EDURE	
	B is	added to A, then C is added to the	mixture and
	then	D is added to the mixture.	
10			
	Exam	ple 16: Massage care gel	
	FORM	ULA	
	Α	Compound of example 3:	3.00%
		Water:	30%
15			
	В	Sepicide [™] CI:	0.20%
		Sepicide [™] HB:	0.30%
		Fragrance:	0.05%
20	С	Colorant:	q.s.
		Water: q.s. for	100%
	D	Micropearl™ SQL:	5.0%
		Lanol [™] 1688:	2%
25			
23	PROCI	EDURE	
		prepared, B is added, then C is added	and then D
		dded.	
	ID a	adea.	
30	Exam	ple 17: Radiance gel	
	FORM	- ULA	
	A	Compound from example 4:	4%
		Water:	30%
35	В	Elastine HPM:	5.0%
	C	Micropearl™ M 100:	3 %
		Water:	5%

5	A is	Sepicide [™] CI: Sepicide [™] HB: Fragrance: 50% sodium pyrrolidinonecarbon Water: EDURE prepared, B is added, then Codded.	q.s.	for	
10	15 4	aaca.			
10	Exam	ple 18 : Body milk			
	FORM				
	A	—— Sepiperl™ N:			3.0%
		Glyceryl triheptonate:		10.09	ę .
15					
	В	Water:	q.s.	for	100%
	С	Compound from example 5:			1.0%
20	D	Eragrango			a a
20	ט	Fragrance: Preservative:			q.s. q.s.
		rieservative.			4.5.
25	A is	EDURE melted at approximately 75°C. 5°C, then C is added at approx added.			
	Exam	ple 19: Make-up-removing emula	sion c	compr	ising sweet
	almo	nd oil			_
30	FORM	ULA	•		
		Montanov [™] 68:			5%
		Sweet almond oil:			5%
		Water:	q.s.	for	100%
		Compound of example 4:			0.3%
35		Glycerol:			5%

Preservative:

Fragrance:

0.2%

0.3%

Example 20: Moisturizing cream for greasy skin FORMULA

	Montanov [™] 68:		5%
	Cetylstearyl octanoate:		8%
5	Octyl palmitate:		2%
	Water:	q.s. for	100%
	Compound of example 3:		0.6%
	Micropearl™ M100:		3.0%
	Mucopolysaccharides:		5%
10	Sepicide™ HB:		0.8%
	Fragrance:		0.3%

Example 21: Alcohol-free soothing aftershave balm FORMULA

15	Mixture of lauryl amino a	acids:	0.1% to 5%
	Magnesium potassium aspan	ctate:	0.002% to 0.5%
	Lanol™ 99:	•	2%
	Sweet almond oil:		0.5%
	Water:	q.s. for	100%
20	Compound of example 2:		3%
	Sepicide™ HB:		0.3%
	Sepicide [™] CI:	0.2%	
	Fragrance:		0.4%

25 Example 22: Cream with AHAs for sensitive skin FORMULA

	Mixture of lauryl amino acids:	0.1% to 5%
	Magnesium potassium aspartate:	0.002% to 0.5%
	Lanol [™] 99:	2%
30	Montanov [™] 68:	5.0%
	Water:	q.s. for 100%
	Compound of example 2:	1.50%
	Gluconic acid:	1.50%
	Triethanolamine:	0.9%
35	Sepicide™ HB:	0.3%
	Sepicide™ CI:	0.2%
	Fragrance:	0.4%

Example 23: Aftersum soothing care preparation

	FORMULA	
	Mixture of lauryl amino acids:	0.1% to 5%
	Magnesium potassium aspartate:	0.002% to 0.5%
5	Lanol™ 99:	10.0%
	Water: q.s. for	r 100%
	Compound of example 4:	2.50%
	Sepicide™ HB:	0.3%
	Sepicide™ CI:	0.2%
10	Fragrance:	0.4%
	Colorant:	0.03%
	Example 24: Make-up-removing milk	
	FORMULA	
15	Sepiperl TM N:	3%
	Primol™ 352:	8.0%
	Sweet almond oil:	2%
	Water: q.s. for	r 100%
	Compound of example 3:	0.8%
20	Preservative:	0.2%
	Example 25: Body milk	
	FORMULA	
	Sepiperl™ N:	3.5%
25	Lanol™ 37T:	8.0%
	${ t Solagum}^{ t TM}$ L:	0.05%
	Water: q.s	s. for 100%
	Benzophenone:	2.0%
	Dimethicone 350 cPs:	0.05%
30	Compound of example 5:	0.8%
	Preservative:	0.2%
	Fragrance:	0.4%
	Example 26: Fluid emulsion with an alk	kaline pH
35	Marcol™ 82:	5.0%
	NaOH:	10.0%
	Water: q.s	s. for 100%
	Compound of example 2:	1.5%

Example 27: Liquid foundation

	Example 27: Liquid foundation	
	FORMULA	
	Simulsol™ 165:	5.0%
	$Lanol^{TM}$ 84D:	8.0%
5	Lanol TM 99:	5.0%
	Water: q.s. for	100%
	Inorganic pigments and fillers:	10.0%
	Compound of example 3:	1.2%
	Preservative:	0.2%
10	Fragrance:	0.4%
	Example 28: Antisun milk	
	FORMULA	
	Sepiperl™ N:	3.5%
15	Lanol™ 37T:	10.0%
	Parsol™ NOX:	5.0%
	Eusolex [™] 4360:	2.0%
	Water: q.s. for	100%
	Compound of example 4:	1.8%
20	Preservative:	0.2%
	Fragrance:	0.4%
	Example 29: Eye contour gel	
	FORMULA	
25	Compound of example 3:	2.0%
	Fragrance:	0.06%
	Sodium pyrrolidinonecarboxylate:	0.2%
	Dow Corning [™] 245 Fluid:	2.0%
	Water: q.s. for	100%
30		
	Example 30: Leave-on care composition	
	FORMULA	
	Compound of example 4:	1.5%
	Fragrance:	q.s.
35	Preservative:	q.s.
	Dow Corning™ X2 8360:	5.0%
	Dow Corning [™] Q2 1401:	15.0%
	Water: q.s. for	100%

	Exam	ple 31: Slimming gel		
		Compound of example 5:		5%
		Ethanol:		30%
		Menthol:		0.1%
5		Caffeine:		2.5%
		Ruscus extract:		2%
		Ivy extract:		2%
		Sepicide™ HB:		1%
		Water:	q.s. for	100%
10				
	Exam	ple 32: Alcohol-free soothing	aftershave	balm
	FORM	<u>ULA</u>		
	Α	Lipacide™ PVB:		1.0%
		Lanol™ 99:		2.0%
15		Sweet almond oil:		0.5%
	В	Compound of example 3:		3.5%
	C	Water:	q.s. for	100%
20				
	D	Fragrance:		0.4%
		Sepicide [™] HB:		0.4%
		Sepicide™ CI:		0.2%
25	Exam	ple 33: Aftershave refreshing	gel	
		FORMULA		0
	A	Lipacide™ PVB:		0.5%
		Lanol™ 99:		5.0%
		Compound of example 2:		2.5%
30				
	В	Water:	q.s. for	100%
	a	7. TM T.		0 50
	С	Micropearl™ LM:		0.5%
2.5		Fragrance:		0.2%
35		Sepicide [™] HB:		0.3%
		Sepicide [™] CI:		0.2%

	- 35 -			
Exar	nple 34: Care preparation for	greasy	skin	
FORM	ALUN			
Α	Micropearl™ M310:			1.0%
	Compound of example 4:			5.0%
	Octyl isononanoate:			4.0%
В	Water:	q.s.	for	100%
С	Sepicontrol [™] A5:			4.0%
	Fragrance:			0.1%
	Sepicide™ HB:			0.3%
	Sepicide [™] CI:			0.2%
D	Capigel™ 98:			0.5%
	Water:			10%
Exar	mple 35: Cream comprising AHAs			
FORM	MULA			
A	Montanov™ 68:			5.0%
	Lipacide™ PVB:			1.05%
	Lanol™ 99:			10.0%
В	Water:	q.s.	for	100%
	Gluconic acid			1.5%
	TEA (triethanolamine):			0.9%
С	Compound of example 5:			1.5%

Example 36: Non-greasy self-tanning preparation for the face and body

0.4%

0.2%

0.4%

35 FORMULA

D

Fragrance:

Sepicide TM HB:

 ${\tt Sepicide^{TM}}\ {\tt CI:}$

5

10

15

20

25

30

A	Lanol™ 2681:	3.0%
	Compound of example 4:	2.5%

B Water: q.s. for 100%

		Dihydroxyacetone:		3.0%
	С	Fragrance: Sepicide™ HB:		0.2%
5		Sodium hydroxide:	q.s. pH :	
	Exan	mple 37: Antisun milk compris	sing Tahit	ian perfumed
		MULA		
10	A	—— Tahitian perfumed oil		10%
		Lipacide™ PVB:		0.5%
		Compound of example 2:		2.2%
	В	Water:	q.s. for	100%
15				
	С	Fragrance:		0.1%
		Sepicide™ HB:		0.3%
		Sepicide [™] CI:		0.1%
		Octyl methoxycinnamate:		4.0%
Example 38: Antisun care preparation for the face FORMULA				
	Α	Cyclomethicone and dimethicon	nol:	4.0%
		Compound of example 3:		3.5%
25				
	В	Water:	q.s. for	100%
	С	Fragrance:		0.1%
		Sepicide [™] HB:		0.3%
30		Sepicide™ CI:		0.21%
		Octyl methoxycinnamate:		5.0%
		Titanium oxide-coated mica		2.0%
		Lactic acid:	q.s. for	pH = 6.5
35	-			
		MULA		3.50
	A	Lanol TM 99:		15%
		Montanov [™] 68:		5.0%
		Octyl para-methoxycinnamate:		3.0%

	В	Water:	q.s.	for	100%
		Dihydroxyacetone:			5.0%
		Monosodium phosphate:			0.2%
5					
	C	Compound of example 4:			0.5%
	D	Fragrance:			0.3%
		Sepicide™ HB:			0.8%
10		Sodium hydroxide:	q.s.	for	pH = 5.
	Examp	ole 40: Sheen gel			
		Compound of example 5:			1.5%
		Volatile silicone:			25%
15		Monopropylene glycol:			25%
		Demineralized water:			10%
		Glycerol:	q.s.	for	100%
	Examp	ole 41: Slimming gel			
20	_	Compound of example 4:			1.5%
		Isononyl isononanoate:			2%
		Caffeine:			5%
		Ethanol:			40%
		Micropearl TM LM:			2%
25		Demineralized water:	q.s.	for	100% .
		Preservative, fragrance:			q.s.
	Exam	ole 42: Make-up-removing milk			
	•	Simulsol TM 165:			4%
30		Montanov TM 202:			1%
		Caprylate/caprate triglyceride	e:		15%
		Pecosil™ DCT:			1%
		Demineralized water:			q.s.
		Capigel [™] 98:			0.5%
35		Compound of example 5			1%
		Proteol™ OAT:			2%
		Sodium hydroxide:	q.s.	for	pH = 7

Example 43: Restructuring rinse-off cream mask for stressed and embrittled hair

	FORMULA	
	Ketrol™ T:	0.5%
5	Pecosil™ SPP50:	0.75%
	N-Cocoyl amino acids	0.70%
	Butylene glycol:	3.0%
	Compound of example 1:	3.0%
	Montanov TM 82:	3.0%
10	Jojoba oil:	1.0%
	Lanol™ P:	6.0%
	Amonyl™ DM:	1.0%
	Lanol™ 99:	5.0%
	Sepicide™ HB:	0.3%
15	Sepicide™ CI:	0.2%
	Fragrance:	0.2%
	Water q.s. for	100%
	Example 44: Antisun cream	
20	Simulsol [™] 165:	3%
	Montanov™ 202:	2%
	C ₁₂ -C ₁₅ benzoate:	8%
	Pecosil [™] PS 100:	2%
	Dimethicone:	2%
25	Cyclomethicone:	5%
	Octyl para-methoxycinnamate:	6%
	Benzophenone-3:	4%
	Titanium oxide	8%
	Xanthan gum:	0.2%
30	Butylene glycol:	5%
	Demineralized water: q.s. for	100%
	Compound of example 2:	1.5%
	Preservative, fragrance:	q.s.
35	Example 45: Care gel for combination skin	
	Compound of example 3:	48
	Vegetable squalane:	5%
	Dimethicone:	1.5%

4%

Sepicontrol[™] A5:

	Xanthan gum:			0.3%
	Water:	q.s.	for	100%
	Preservative, fragrance:			
5	Example 46: Hair lotion			
	FORMULA			
	Butylene glycol:			3.0%
	Compound of example 6:			3%
	Simulsol™ 1293:			3.0%
10	Lactic acid:	q.s.	for	pH = 6
	Sepicide™ HB:			0.2%
	Sepicide [™] CI:			0.3%
	Fragrance:			0.3%
	Water:		q.s.	100%
15				
	Example 47: Protecting and relaxi	ng sha	mpoo	
	FORMULA			
	Amonyl™ 675 SB:			5.0%
	28% Sodium lauryl ether sulf	ate:		35.0%
20	Composition of example 6:			3.0%
	Sepicide™ HB:			0.5%
	Sepicide $^{\mathtt{TM}}$ CI:			0.3%
	Sodium hydroxide:		q.s.	pH = 7.2
	Fragrance:			0.3%
25	Colorant (FDC Blue 1/Yellow	5):		q.s.
	Water:	q.s.	for	100%
	Example 48: Leave-on protecting	prepar	ation	; Hair care
	preparation for combating stress			
30	FORMULA			
	Ketrol™ T:	_		0.5%
	Mixture of cocoyl amino acid	ds:		3.0%
	Butylene glycol:			5.0%
	DC 1501:			5.0%
35	Composition of example 1:			4.0%
	Sepicide™ HB:			0.5%
	Sepicide [™] CI:			0.3%
	Fragrance:			0.3%
	Water:	q.s.	for	100%

Example 49: Vitamin cream

•	Simulsol™ 165:	5%
	Montanov [™] 202:	1%
5	Caprylic/capric triglycerides:	20%
	Vitamin A palmitate:	0.2%
	Vitamin E acetate:	1%
	Micropearl TM M 305	1.5%
	Compound of example 1:	2%
10	Water q.s. for	100%
	Preservative, fragrance	q.s.

The definitions of the commercial products used in the examples are as follows:

15 Simulsol[™] 1293 is hydrogenated and ethoxylated castor oil with an ethoxylation number of 40, sold by Seppic.

Capigel[™] 98 is a liquid thickener based on acrylate copolymer sold by Seppic.

 $Ketrol^{TM}$ 99 is xanthan gum, sold by Kelco.

20 Lanol[™] 99 is isononyl isononanoate, sold by Seppic.

DC1501 is a mixture of cyclopentasiloxane and dimethiconol sold by Dow Chemical.

 $Montanov^{TM}$ 82 is an emulsifying agent based on cetearyl alcohol and on cocoyl glucoside.

25 Montanov[™] 68 (cetearyl glucoside) is a selfemulsifiable composition, such as disclosed in WO 92/06778, sold by Seppic.

 $Micropearl^{TM}$ M 100 is an ultrafine powder with a very soft feel and with a mattifying action, sold by

30 Matsumo.

SepicideTM CI, imidazolidineurea, is a preservative sold by Seppic.

Pemulen[™] TR is an acrylic polymer sold by Goodrich.

 $Simulsol^{TM}$ 165 is self-emulsifiable glyceryl stearate,

35 sold by Seppic.

Lanol $^{\text{TM}}$ 1688 is an emollient ester with a non-greasy effect sold by Seppic.

 $Lanol^{TM}$ 14M and $Lanol^{TM}$ S are consistency factors sold by Seppic.

Sepicide[™] HB, which is a mixture of phenoxyethanol, methylparaben, ethylparaben, propylparaben and butylparaben, is a preservative sold by Seppic.

Monteine™ CA is a moisturizing agent sold by Seppic.

5 SchercemolTM OP is an emollient ester with a non-greasy effect.

Lanol $^{\text{TM}}$ P is an additive with a stabilizing effect sold by Seppic.

Parsol[™] MCX is octyl para-methoxycinnamate, sold by Givaudan.

Sepiperl $^{\text{TM}}$ N is a pearlescent agent, sold by Seppic, based on a mixture of alkyl polyglucosides such as those disclosed in WO 95/13863.

Micropearl[™] SQL is a mixture of microparticles including squalane which is released under the action of massaging; it is sold by Matsumo.

Lanol[™] 99 is isononyl isononanoate, sold by Seppic.

 $Lanol^{TM}$ 37T is glyceryl triheptanoate, sold by Seppic.

 $Solagum^{TM}$ L is a carrageenan sold by Seppic.

20 Marcol[™] 82 is a liquid paraffin sold by Exxon. Lanol[™] 84D is dioctyl malate, sold by Seppic.

Parsol[™] NOX is a sunscreen sold by Givaudan.

EusolexTM 4360 is a sunscreen sold by Merck.

Dow CorningTM 245 Fluid is cyclomethicone, sold by Dow

25 Corning.

 $Lipacide^{TM}$ PVB is a hydrolyzate of acylated wheat proteins sold by Seppic.

Micropearl $^{\text{TM}}$ LM is a mixture of squalane, poly(methyl methacrylate) and menthol sold by Seppic.

30 Sepicontrol[™] A5 is a capryloylglycine, sarcosine, extract of Cinnamon zylanicum mixture sold by Seppic, such as those disclosed in international patent application PCT/FR98/01313 filed on 23 June 1998.

 $Lanol^{TM}$ 2681 is a coconut caprylate/caprate mixture sold

35 by Seppic.

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Montanov $^{\text{TM}}$ 202 is an APG/fatty alcohols composition as disclosed in WO 98/47610, sold by Seppic.